Haplotypes at the *miR156* and *miR172* transcript regions show a significant response to divergent recurrent selection for vegetative phase change in maize

Jessica Rutkoski, Eric Riedeman, William Tracy, and Shawn Kaepppler.

**Abstract**

Sweet corn (*Zea mays* L.) plants that transition from juvenility to the adult phase earlier during development are generally more resistant to common rust (*Puccinia sorghi*), a disease that causes substantial yield loss in sweet corn. This developmental transition is called vegetative phase change (VPC). VPC is regulated in part by *miR156* and *miR172*. We found that natural variation at the *miR156* and *miR172* transcript regions are important in determining variation in the timing of VPC. We sequenced the *miR156* and *miR172* transcript regions in populations originating from the same source population and divergently selected for eight generations for early and late VPC. For both the *miR156* and *miR172* transcript regions we detected four distinct haplotypes. Overall and individual haplotype frequencies showed significant responses to selection for both early and late VPC. These data contribute to the identification of important loci involved in VPC. Understanding the genetic basis for VPC will facilitate maize breeding aimed at combining early VPC with other desirable traits.

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Abstract

Sweet corn (Zea mays L.) plants that transition from juvenility to the adult phase earlier during development are generally more resistant to common rust (Puccinia sorghi), a disease that causes substantial yield loss in sweet corn. This developmental transition is called vegetative phase change (VPC). VPC is regulated in part by miR156 and miR172. We found that natural variation at the miR156 and miR172 transcript regions are important in determining variation in the timing of VPC. We sequenced the miR156 and miR172 transcript regions in populations originating from the same source population and divergently selected for eight generations for early and late VPC. For both the miR156 and miR172 transcript regions we detected four distinct haplotypes. Overall and individual haplotype frequencies showed significant responses to selection for both early and late VPC. These data contribute to the identification of important loci involved in VPC. Understanding the genetic basis for VPC will facilitate maize breeding aimed at combining early VPC with other desirable traits.

Introduction:

Common rust is the most prevalent disease afflicting sweet corn (Zea mays L.) grown in the upper Midwest. Rust is caused by the fungal pathogen Puccinia sorghi, which is often controlled by applying fungicides to crops and by growing sweet corn with the Rp1-D allele conferring genetic resistance. Since 1999, new races of Puccinia sorghi have appeared in the Midwest that are able to infect sweet corn plants with the Rp1-D allele making this resistance gene ineffective (Pataky et al. 2001). This study investigates a new form of genetic resistance
that exploits partial adult plant resistance to rust that occurs in adult sweet corn plants. The two vegetative phases in maize, juvenile and adult, are distinct in that adult leaves have a thicker cuticle and juvenile leaves have more epicuticular wax (Poethig, 1990). Juvenile leaves are more susceptible to *P. sorghi*. Plants with early or late vegetative phase change can be selected to decrease or increase the duration of the juvenile phase; those with a longer juvenile phase experience greater susceptibility to common rust and insects (Basso et al., 2008; Abedon and Tracy, 1996).

Vegetative phase change (VPC) in maize is regulated in part by *microRNA156* (*miR156*) and *microRNA172* (*miR172*). MicroRNAs are small RNA sequences that cleave complementary mRNA sequences, therefore reducing the level of the protein product the target mRNA codes for. Overexpression of *miR156* increases the duration of the juvenile phase (Chuck et al., 2007). *miR156* cleaves target mRNA that code for specific transcription factors that promote transcription of developmental genes. The transcription factors are therefore less numerous and less of the corresponding developmental gene is expressed. (Rhodes et al. 2002). *gl15* is an essential gene for juvenile leaf trait expression. *miR172* down regulates the juvenility gene *gl15*.
by cleaving the transcript of \textit{gl15}. This promotes the transition to the adult vegetative phase (Lauter et al., 2005).

From an original Minn 11 sugary1 sweet corn population, Professor Bill Tracy has used divergent recurrent selection selecting VPC in the late and early direction for eight cycles in an ongoing study. The timing of VPC has been very responsive. (Figure 1) (Riedeman et al., 2008). Morphological changes have resulted and susceptibility to rust has increased in the cycles selected in the late phase change direction. In this experiment, we used sequencing to test the hypothesis that the haplotype frequencies at the \textit{miR156} and \textit{miR172} transcript regions were altered in response to selection and whether the natural variation at the \textit{miR156} transcript region and the \textit{miR172} transcript region is important in determining the observed variation of the timing of vegetative phase change (VPC). Understanding the genetic basis for regulation of VPC in maize will allow breeders to produce maize that combines high quality traits with early VPC, conferring partial yet durable resistance to rust (Basso et al., 2008, Abedon and Tracy, 1996).

**Materials and Methods**

**Population Development:** Minn 11, a sugary1 sweet corn population, formed cycle 0 (C0) of a divergent recurrent selection program. While the parents of Minn 11 included both Corn Belt Dent and sweet corn germplasm, the exact parents are unknown. Approximately 150 C0 plants were self-pollinated, harvested, and planted in family rows. Family rows were evaluated for the average last leaf with juvenile wax. The earliest and latest 20\% for last leaf with juvenile wax intermated using bulked pollen from the selections. A minimum of 100 ears was obtained for each direction of selection. Equal amounts of seed from these ears were pooled to form balance bulks for each direction. Remnant seed was planted ear to row for the next cycle of selection. This process was repeated through eight cycles of selection in both directions. The original
population (C0), cycle 8 in the early direction of selection (C8E), and cycle 8 in the late direction of selection (C8L) were the populations evaluated for haplotype frequency in this experiment.

**Leaf harvest and preparation:** We grew 120 plants from C0, and the C8E and C8L. Immature leaves were harvested, freeze-dried and pulverized for DNA extraction.

**DNA extraction:** We isolated DNA using the DNA isolation using CTAB protocol (Kidwell, K.K., and T.C. Osborn, 1993). We measured DNA concentration using a nano-drop spectrophotometer.

**Population parentage testing:** Four potential parents of the C0 population, B73, Pa405, Ia453, and Ii125b, were screened using microsatellite (SSR) markers. DNA from the four potential parents and 96 individuals from C0 were PCR amplified with random SSR marker primers. Polymorphisms were screened using the Mega-Gel protocol for polyacrylamide gel electrophoresis. Each gel mix contained 75 ml 1x TBE, 22.5 ml 40% acrylamide stock 1.05 ml 10% APS, 120 µl TEMED, and 51 ml of ddH2O. PCR amplified samples were stained using Type IV loading dye (6x) and 1.25µl of a 1/100 dilution of Gel Red per 10µl. Stained samples were run on a gel at 7 watts for 2 hours, a second set of samples was run then loaded and run for 2 hours. Banding patterns on the gel were visualized and photographed using a UV light box.

**DNA amplification:** Three bulks of DNA, 100ng from each individual within its respective population, were made one each for C0, C8L, and C8E. We used and optimized PCR protocol to amplify each DNA bulk. We used a GoTaq® Hot Start Polymerase kit from Promega. Our reaction mixes contained 2µl of 10x colorless buffer, 0.4µl dNTP, 0.65µl forward and reverse primers, 0.1µl HotStar Taq, 0.8ng dry DNA and 16.05µl ddH2O, total reaction amount was 20ul. We used the miR156 forward primer GTGGCGTCAACAACATTACG, and reverse primer AAGGCATAACCAACAATCA which flanks a region about 1900 bps downstream from the
miR156 transcript region. We used the miR172 forward primer ACAAAGGCCAAGCGCTACTA, and reverse primer CACATGGGTGACGATGCTAC which flanks a region about 1500 bps downstream of the miR172 transcript region. We amplified each bulk once using primers for the miR156 transcript region and again for the miR172 transcript region (six PCR products total). Specific priming and amplification of these regions was confirmed using gel electrophoresis on 1% agarose gels. Resolution of more than one band on the gel indicated non-specific priming or DNA degradation.

**DNA purification:** We cut the primer specific bands out of the 1% agarose gel to ensure isolation of primer specific DNA. Used a QIAquick spin column kit to remove primers, nucleotides, enzymes, mineral salts, agarose, ethidium bromide, and other impurities from the DNA samples. Purified DNA was concentrated with EtOH DNA precipitation, and resuspended in 10ul TE.

**Cloning into bacteria:** We ligated each of the six purified PCR products into a pGEM-T Easy Vector using T4 DNA ligase. Ligation of control insert DNA into the vector was used as a positive control and ligation of the empty vector without insert DNA was used as a negative control. We prepared 2 LB/ampicillin/IPTG/X-Gal plates for each ligation reaction and equilibrated the plates to room temperature prior to plating. We transformed JM109 High Efficiency Competent Cells by mixing 2ul of each ligation reaction with 50ul of competent cells. The mix was placed on ice for 20 minutes, heat-shocked for 45-50 seconds in a 42°C water bath, and returned to ice for 2 minutes. We incubated the transformed cells for 1.5 hours at 37°C with shaking and plated 100ul of each culture onto duplicate LB/ampicillin/IPTG/X-Gal plates. Plates were incubated overnight at 37°C. White colonies contained inserts. Individual white colonies
were transferred to a replicate plate and a well in a 96 well PCR plate. One plate was designated for each of the 6 transformation reactions.

**Clean up and Sequencing:** We PCR amplified DNA from transformed colonies. PCR products were diluted 2:40 with TE. We prepared our each of the six 96 well plates for dye-terminator sequencing using the BigDye terminator kit containing fluorescently labeled dideoxy terminators. Only the plate’s corresponding forward primers were used. The reactions were run on a thermocycler using the BigDye protocol. DNA was cleaned up and sequenced at the UW Madison Biotechnology DNA Sequence Laboratory.

**Analysis:** DNA sequences were aligned using BioEdit software and significance in changes of haplotype frequency was determined using SAS. We ran two-way chi-square tests when over 80% of expected values were over 5. Fishers exact tests used when this criteria was not met. The significance level for both tests was set at $\alpha= 0.05$.

**Results:**

**Population parentage:**

We aimed to identify the parents of C0 in order to develop SNP markers that could be used to screen C0, C8E, and C8L for haplotype frequency changes. We found that SSR Markers Bnlg-1144 and Umc-1014 produced bands present in C0 that were absent in the potential parents. We thus eliminated B73, Pa405, Ia453, and Il125b as the parents of C0 (Figure 2). We therefore proceeded to detect haplotype frequency changes between C0, C8E, and C8L via sequencing.
Haplotype Frequency Analysis:

From sequence data at the miR156 transcript region six single nucleotide polymorphisms (SNPs) were found. Five of the SNPs were organized into four distinct haplotypes. Overall haplotype frequencies varied significantly between C0 and C8E as well as between the C8L and C8E populations (Table 1). Using a two-way chi-square test, we found that haplotype frequencies between C8E and C8L were significantly different (p=0.0003). Using a fisher’s exact test we found that haplotype frequencies were significantly different between the C0 and C8E (p=0.0026), but were not significantly different between C0 and C8L (p=0.3028). These data suggest that the miR156 transcript region is a target for selection for early VPC.

From sequence data at the miR172 transcript region eight SNPs and nine insertion/deletion polymorphisms were found. All these polymorphisms were organized into four distinct haplotypes. Overall haplotype frequencies varied significantly between C0 and C8L as well as between the C8L and C8E populations (Table 1). Using a two-way chi-square test, we found that haplotype frequencies were significantly different between C8E and the C8L (p=0.0002), but were not significantly different between C0 and the C8E (p=0.1013). Using a fisher’s exact test, we found that haplotype frequencies were significantly different between C0 and the C8L (p=0.0231). These data suggest that the miR172 transcript region is a target for selection for late VPC.

Figure 2: SSR markers umc-1014 (left) and bnlg-1144 (right) resolve bands in the C0 individuals that are not present in the potential parents, B73, Pa405, Ia453, and II125b. These candidates are thus eliminated as the parents of C0.
### Comparison pairs

<table>
<thead>
<tr>
<th>Comparison pairs</th>
<th><em>miR156</em> transcript region</th>
<th><em>miR172</em> transcript region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cycle 8 early vs. Cycle 0</td>
<td>p=0.0026</td>
<td>p=0.1013</td>
</tr>
<tr>
<td>Cycle 8 late vs. Cycle 0</td>
<td>p=0.3028</td>
<td>p=0.0231</td>
</tr>
<tr>
<td>Cycle 8 late vs. Cycle 8 early</td>
<td>p=0.0003</td>
<td>p=0.0002</td>
</tr>
</tbody>
</table>

Table 1: Statistical significance of overall haplotype frequency among all possible comparisons of C0, C8E and C8L

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**Figure 3:** Haplotype frequencies at the *miR156* transcript region for C80, C8E, and C8L

![miR156 haplotype frequencies](chart1)

<table>
<thead>
<tr>
<th>miR156 haplotype frequencies in C0, C8E and C8L</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haplotype 1</td>
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<tr>
<td>haplotype 1</td>
</tr>
<tr>
<td>haplotype 2</td>
</tr>
<tr>
<td>haplotype 3</td>
</tr>
<tr>
<td>haplotype 4</td>
</tr>
</tbody>
</table>

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**Figure 4:** Haplotype frequencies at the *miR172* transcript region for C80, C8E, and C8L

![miR172 haplotype frequencies](chart2)

<table>
<thead>
<tr>
<th>miR172 haplotype frequencies in C0, C8E and C8L</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haplotype 1</td>
</tr>
<tr>
<td>haplotype 1</td>
</tr>
<tr>
<td>haplotype 2</td>
</tr>
<tr>
<td>haplotype 3</td>
</tr>
<tr>
<td>haplotype 4</td>
</tr>
</tbody>
</table>
Frequencies of individual haplotypes at the *miR156* and *miR172* transcript regions varied between C0, C8E, and C8L (Figure 3, Figure 4). Using two-way chi-square tests, we found that the frequency of haplotype one at the *miR156* transcript region was significantly increased by selection for early VPC (p=0.0105) and was significantly different between C8E and C8L. The frequency of haplotype four at the *miR156* transcript region was significantly decreased by selection for early VPC (p=0.096) and was significantly different between C8E and C8L (Table 2).

<table>
<thead>
<tr>
<th>Comparison pairs</th>
<th>Haplotype 1</th>
<th>Haplotype 2</th>
<th>Haplotype 3</th>
<th>Haplotype 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cycle 8 early vs. Cycle 0</td>
<td>0.0105</td>
<td>0.9634</td>
<td>0.6389</td>
<td>0.0096</td>
</tr>
<tr>
<td>Cycle 8 late vs. Cycle 0</td>
<td>0.7254</td>
<td>0.1242</td>
<td>0.329</td>
<td>0.9483</td>
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<td>Cycle 8 late vs. Cycle 8 early</td>
<td>0.0034</td>
<td>0.1227</td>
<td>0.1571</td>
<td>0.007</td>
</tr>
</tbody>
</table>

Table 2: Statistical significance of individual haplotype frequencies at the *miR156* transcript region among all possible comparisons of C0, C8E, and C8L

Using two-way chi-square tests we found that the frequency of haplotype three at the *miR172* transcript region was significantly decreased by selection for late VPC (p=0.0167) and was significantly different between C8L and C8E (p=0.0018). The frequency of haplotype four at the *miR172* transcript region was significantly different between C8E and C8L (p=0.000669) (Table 3).

<table>
<thead>
<tr>
<th>Comparison pairs</th>
<th>Haplotype 1</th>
<th>Haplotype 2</th>
<th>Haplotype 3</th>
<th>Haplotype 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cycle 8 early vs. Cycle 0</td>
<td>0.4633</td>
<td>0.1145</td>
<td>0.5347</td>
<td>0.0592</td>
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<tr>
<td>Cycle 8 late vs. Cycle 0</td>
<td>0.8034</td>
<td>0.3597</td>
<td>0.0167</td>
<td>0.1209</td>
</tr>
<tr>
<td>Cycle 8 late vs. Cycle 8 early</td>
<td>0.3332</td>
<td>0.5177</td>
<td>0.0018</td>
<td>0.000669</td>
</tr>
</tbody>
</table>

Table 3: Statistical significance of individual haplotype frequencies at the *miR172* transcript region among all possible comparisons of C0, C8E, and C8L
**Discussion:**

Our analysis of sequence data 1900bps downstream from the miR156 and 1500bps downstream from the miR172 transcript regions reveals sequence variation within both regions. The variation in each region is organized into four distinct haplotypes that show a non-random response to selection for VPC. Haplotypes at the miR172 transcript region significantly respond to selection for late VPC whereas haplotypes at the miR156 transcript region significantly respond to selection for early VPC. These haplotypes may be linked to targets for selection in the pre-miRNA sequences or enhancer regions. These presumed targets may affect mature miRNA levels through various mechanisms. Potential mechanisms include: reduced DNA transcription at the miR156 and miR172 transcript regions, or reduced stability of pre-miRNA156 and pre-miRNA172 secondary structures. Although microRNA sequences are highly conserved, within species pre-miRNA polymorphisms have been found within populations of *Arabidopsis thaliana*. In *A. thaliana* these pre-miRNA polymorphisms alter pre-miRNA secondary structure and can reduce their stability thus reducing mature microRNA levels. (de Meaux et. al., 2008, Ehrenreich et. al., 2008). Polymorphisms in the pre-miR156 and pre-miR172 sequence may exist in maize and may be targets for selection linked to the haplotypes detected in this experiment. Interestingly, if selection reduced levels of miR156 early VPC would be promoted while reduced levels of miR172 would promote late VPC.

**Conclusion:**

**Population Parentage:**

Our parentage analysis of C0 did not succeed in identifying the parents of the population. Had we identified the parents we would have sequenced the parents at the miR156 and miR172 transcript regions to develop SNP markers. These SNP markers would have been used to screen
C0, C8E, and C8L for haplotype frequency changes between populations. With no parents identified, we detected the haplotype frequency changes using sequencing. Developing SNP markers from the C0 sequence data at the *miR156* and *miR172* transcript region would be useful to facilitate further studies of haplotype frequency changes at these regions in populations derived from C0.

**Haplotype Frequency Analysis:**

Our observation that haplotype frequencies at the *miR156* and *miR172* transcript regions significantly respond to selection for the timing of VPC supports the hypothesis that the variation at these regions is involved in the differential timing of VPC observed in C0, C8E, and C8L. These data contribute to the identification of important loci involved in VPC.

To validate the importance of variation at the *miR156* and *miR172* transcript regions in the response to selection for VPC intermediate cycles of selection should be haplotyped to determine whether there was a consistent response to selection. To investigate the genetic mechanism employed by the targets for selection in this experiment MIR156 and MIR172 levels in C0, C8E and C8L populations should be tested. If MIR156 and MIR172 levels differ, the stability of the pre-miRNA156 and pre-miRNA172 in should be investigated.

Understanding the genetic basis for VPC will facilitate maize breeding aimed at combining either early or late VPC with other desirable traits. Since early VPC in maize imparts partial resistance to common rust, incorporating early VPC into maize cultivars could reduce crop losses in Wisconsin where rust is prevalent.
References:


Riedeman, E.S., M.A. Chandler, and W.F. Tracy. 2008. Seven cycles of divergent recurrent selection for vegetative phase change and indirect effects on resistance to common rust (*Puccinia sorghi*) and European corn borer (*Ostrinia nubilalis*). Accepted by Crop Science.
